

DECLARATION OF CONFORMITY TO Regulation(EU) 2017/745 CONCERNING MEDICAL DEVICES

MANUFACTURER: Edan Instruments, Inc.
#15 Jinhui Road, Jinsha Community, Kengzi Sub-District,
Pingshan District, 518122 Shenzhen, P.R.China
SRN: CN-MF-000009957

EUROPEAN REPRESENTATIVE: Shanghai International Holding Corp. GmbH (Europe)
Eiffestrasse 80, 20537 Hamburg Germany
SRN: DE-AR-000000001

PRODUCT/MODEL: Ultrasonic Pocket Doppler/SONOTRAX Lite,
SONOTRAX Basic, SONOTRAX Pro, SONOTRAX Basic
A, SONOTRAX II, SONOTRAX II Pro

EMDN [NAME/CODE]: FOETAL HEARTBEAT DETECTORS / Z12080103
Basic UDI-DI: 69444138SonotraxSC8
Classification: Class IIa, Rule 10 According To Annex VIII of the MDR
CONFORMITY ASSESSMENT ROUTE: ANNEX IX CHAPTERS I.

WE, EDAN INSTRUMENTS, INC., HERE WITH DECLARE THAT THE ABOVE MENTIONED PRODUCT(S)
MEET THE PROVISIONS OF REGULATION (EU) 2017/745 OF THE EUROPEAN PARLIAMENT AND
OF THE COUNCIL ON MEDICAL DEVICE.

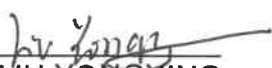
ALL SUPPORTING DOCUMENTATION IS RETAINED AT THE PREMISES OF THE MANUFACTURER.
EU DECLARATION OF CONFORMITY IS ISSUED UNDER THE SOLE RESPONSIBILITY OF THE
MANUFACTURER

STANDARDS APPLIED: IEC 60601-1: 2005+ A1: 2012+ A2: 2020, EN 60601-1-
2:2015+A1:2021, EN 60601-1-6:2010+A2:2021, EN 60601-2-37:2015, EN 61266:
1995, EN ISO 10993-1:2020, EN ISO 10993-5:2009, EN ISO 10993-10:2023, EN ISO
10993-23:2021, EN ISO 780:2015, EN 62304:2006+A1:2015, EN 62366-1: 2015+ A1:
2020, EN ISO 20417: 2021, EN ISO 14971:2019, EN ISO 15223-1:2021

NOTIFIED BODY: TÜV SÜD PRODUCT SERVICE GMBH
RIDLERSTR 65, D-80339 MÜNCHEN, GERMANY

IDENTIFICATION NUMBER 0123
(EC) CERTIFICATE(S): G15 091264 0085 REV.00 VALID UNTIL: 2031-02-17

START OF CE-MARKING: 2002-09-25
PLACE, DATE OF ISSUE: SHENZHEN, 2026.2.27

SIGNATURE: 
NAME LIU YONGYING
MANAGEMENT REPRESENTATIVE